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(54) Title: LIGAND SCREENING AND DESIGN BY X-RAY CRYSTALLOGRAPHY

(57) Abstract

X-ray crystallography can be used to screen compounds that are not known ligands of a target biomolecule for their ability to bind the target biomolecule. The method includes obtaining a crystal of a target biomolecule; exposing the target biomolecule crystal to one or more test samples; and obtaining an X-ray crystal diffraction pattern to determine whether a ligand/receptor complex is formed. The target is exposed to the test samples by either co-crystallizing a biomolecule in the presence of one or more test samples or soaking the biomolecule crystal in a solution of one or more test samples. In another embodiment, structural information from ligand/receptor complexes are used to design ligands that bind tighter, that bind more specifically, that have better biological activity or that have better safety profile. A further embodiment of the invention comprises identifying or designing biologically-active moieties by the instant process. In a further embodiment, a biomolecule crystal having an easily accessible active site is formed by co-crystallizing the biomolecule with a degradable ligand and degrading the ligand.

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Ligand Screening and Design by X-ray Crystallography

Technical Field of the Invention

X-ray crystallography is useful for identifying ligands that bind target receptor molecules and for designing ligands with improved biological activity for the target receptor.

Background of the Invention

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X-ray crystallography (crystallography) is an established, well-studied technique that provides what can be best described as a three-dimensional picture of what a molecule looks like in a crystal. Scientists have used crystallography to solve the crystal structures for many biologically important molecules. Many classes of biomolecules can be studied by crystallography, including, but not limited to, proteins, DNA, RNA and viruses. Scientists have even reported the crystal structures of biomolecules that carry ligands within its receptors (a "ligand-receptor complex").

Given a "picture" of a target biomolecule or a ligand-receptor complex, scientists can look for pockets or receptors where biological activity can take place. Then scientists can experimentally or computationally design high-affinity ligands (or drugs) for the receptors. Computational methods have alternatively been used to screen for the binding of small molecules. However, these previous attempts have met with limited success. Several problems plague ligand design by computational methods. Computational methods are based on estimates rather than exact determinations of the binding energies, and rely on simple calculations when compared with the complex interactions that exist within a biomolecule. Moreover, computational models require experimental confirmation which often expose the models as false positives that do not work on the real target.

Moreover, experimental high-affinity ligand design based on a "picture" of the ligand-receptor complex has been limited to biomolecules that already have known ligands. Finally, scientists only recently reported the crystallographic study of interactions between organic solvents and target biomolecules. Allen et al., <u>J. Phys. Chem.</u>, v. 100, pp. 2605-11 (1996). However, these studies are limited to mapping solvent sites rather than

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ligand sites. It would be desirable to directly identify potential ligands, and to obtain detailed information on how the ligand binds and changes in the target biomolecule. In addition, methods for identifying and/or designing ligands which possess biological and /or pharmaceutical activity with respect to a given target molecule would be desireable.

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Brief Summary of the Invention

Crystallography can be used to screen and identify compounds that are not known ligands of a target biomolecule for their ability to bind the target. The method (hereinafter "CrystaLEAD™") comprises obtaining a crystal of a target biomolecule; exposing the target to one or more test samples that are potential ligands of the target; and determining whether a ligand/biomolecule complex is formed. The target is exposed to potential ligands by various methods, including but not limited to, soaking a crystal in a solution of one or more potential ligands or co-crystallizing a biomolecule in the presence of one or more potential ligands.

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In a further embodiment, structural information from the ligand/receptor complexes found are used to design new ligands that bind tighter, bind more specifically, have better biological activity or have better safety profile than known ligands.

In a preferred embodiment, libraries of "shape-diverse" compounds are used to allow direct identification of the ligand-receptor complex even when the ligand is exposed as part of a mixture. This avoids the need for time-consuming de-convolution of a hit from the mixture. Here, three important steps are achieved simultaneously. The calculated electron density function directly reveals the binding event, identifies the bound compound and provides a detailed 3-D structure of the ligand-receptor complex. In one embodiment, once a hit is found, one could screen a number of analogs or derivatives of the hit for tighter binding or better biological activity by traditional screening methods. 25 Another embodiment uses the hit and information about structure of the target to develop analogs or derivatives with tighter binding or better biological activity. In yet another embodiment, the ligand-receptor complex is exposed to additional iterations of potential ligands so that two or more hits can be linked together to make a more potent ligand.

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Brief Description of the Drawings

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Figure 1 illustrates a structure-based drug design where an initial lead compound is found, and then used as a scaffold to carry additional moieties that fit the subsites that surround a major site.

Figure 2 illustrates a fragment linking approach for a biomolecule having two or more adjacent primary pockets.

Figure 3 is an outline of CrystaLEADTM wherein a crystal is soaked in a solution of various potential ligands (I_1 - I_{10}) and an X-ray diffraction dataset is collected and transformed into an electron density map which is inspected for compound binding.

Figure 4 illustrates a typical compound mixture in 2-D and 3-D. The 3-D figures are theoretical 2Fo-Fc electron density maps that represent the "shape" of the molecules.

Figure 5 is a primary sequence of human urokinase.

Figure 6 illustrates how a hit was detected and identified by shape after urokinase was soaked in a solution containing a mixture of potential ligands. Figure 6A is the initial Fo-Fc map. Figure 6B shows how the compound binds at the active site of urokinase. Figure 6C illustrates the active site without a bound ligand when no compound of the mixture has bound.

Figure 7 illustrates a hit for urokinase soaked in a solution containing a mixture of potential ligands. Figure 7A is the initial Fo-Fc map. Figure 7B shows how the compound binds at the active site of urokinase.

Figure 8 illustrates a hit for urokinase soaked in a solution containing a mixture of potential ligands. Figure 8A is the initial Fo-Fc map. Figure 8B shows how the compound binds at the active site of urokinase.

Figure 9 illustrates two additional hits for urokinase soaked in a solution containing a mixture of potential ligands. Figure 9A is the Fo-Fc map for a strong ligand within the mixture. Figure 9B is the Fo-Fc map for a weaker ligand within the mixture. The weaker ligand was detected only after the strong ligand was removed from the mixture.

Figure 10 illustrates the comparative crystal structures between a lead compound found by CrystaLEAD™ and an optimized follow-up compound.

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Figure 11 illustrates hits that were identified for VanX.

Figure 12 illustrates a hit for urokinase.

Figure 13 illustrates the crystal structure of compound 44 with ErmC'.

Figure 14 illustrates the crystal structure of compound 45 with ErmC'.

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Detailed Description of the Invention

CrystaLEAD™ provides an efficient screening method for identifying compounds that will bind to a target biomolecule. Such compounds can serve as leads or scaffolds to design ligands and/or drugs that have improved biological activity for the target. One must note that tighter binding ligands do not necessarily provide better biological activity or make a better drug, although this is the general rule. It is possible for a weaker binding ligand to provide better biological activity due to factors other than tight binding (e.g., selectivity, bioavailability).

Crystallography has been used extensively to view receptor-ligand complexes for structure-based drug design. To view such complexes, known ligands are usually soaked into the target molecule crystal, followed by crystallography of the complex. Sometimes, it is necessary to co-crystallize the ligands with the target molecule to obtain a suitable crystal.

Until now, crystallography has not been implemented to screen potential ligands despite the detailed structural information that it provides. Possible prejudices against screening compounds by crystallography include the belief that the method is too complicated or time consuming, that suitable crystals are difficult to obtain, that available crystals could not tolerate soaking more than one compound (much less mixtures of ten or more compounds), that too much biomolecule would be needed, that it would be too time consuming to routinely mount crystals, and that constantly changing crystals on the x-ray goniometer would be too tedious.

However, currently available technology has overcome many of these perceived barriers. For example, at one time, molecular targets were only obtained from natural sources and were sometimes unsuitable for crystallization due to natural degradation or glycosylation. In addition, the natural concentration was often too low to obtain the amount of highly purified protein necessary for crystallization. With molecular biology,

large amounts of protein may be expressed and purified for crystallization. When necessary, the protein can even be re-engineered to provide different or better crystal forms.

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Further, brilliant light sources (synchrotron radiation) and more sensitive detectors have become readily available so that the time required to collect data has been reduced dramatically from days to hours or even minutes. Furthermore, existing technologies which are less routine at this time, but may become routine soon, allow full data set collections in the order of seconds or even fractions of a second (e.g., Laue diffraction). J. Hajdu et al., Nature, v. 329, pp. 178-81 (1987). Faster computers and more automation software have greatly decreased the time required for data collection and analysis. Finally, the inventors have discovered that it is possible to soak or co-crystallize mixtures of compounds to screen for potential ligands. Thus, as described below, crystallography is now a practical and feasible screening method.

In CrystaLEAD™, ligands for a target molecule having a crystalline form are identified by exposing a library of small molecules, either singly or in mixtures, to the target (e.g. protein, nucleic acid, etc.). Then, one obtains crystallographic data to compare the electron density map of the putative target-ligand complex with the electron density map of the target biomolecule. The electron density map simultaneously provides direct evidence of ligand binding, identification of the bound ligand, and the detailed 3-D structure of the ligand-target complex. Binding may also be monitored by changes in individual reflections within the crystallographic diffraction pattern which are known to be sensitive to ligand binding at the active site. This could serve as a pre-screen but would not be the primary method of choice because it provides less detailed structural information.

By observing changes in the level of ligand electron density or the intensity of certain reflections in the diffraction pattern as a function of ligand concentration either added to the crystal or in co-crystallization, one may also determine the binding affinities of ligands for biomolecules. Binding affinities may also be obtained by competition experiments. Here, the new compound(s) are soaked or co-crystallized with one of a series of diversely-shaped ligands of known binding affinity. If the known ligand appears in the electron density map, the unknown ligands are weaker binders. However, if one of the new compounds is found to compete for the site, it would be the tightest binder. By

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varying the concentration or identity of the known ligand, a binding constant for the CrystaLEAD™ hit may be estimated.

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The number of compounds screened is based upon the desired detection limit, the compound solubility and the amount of organic co-solvent the crystals will tolerate. Exact numbers depend on each crystal. For example, for a typical crystal that tolerates 1% organic co-solvent, the sensitivity limit would be Kd< 1.5mM to screen 10 compounds simultaneously. For 20 compounds, the sensitivity limit would be Kd<0.63mM. However, crystals that tolerate high organic co-solvents (e.g., 40%), can screen up to 50 compounds within a detection limit of Kd< 1.5mM.

In the most general application of CrystaLEADTM, the hit or lead compound is used to determine what compounds should be tested for biological activity in structure-based drug design. Then derivatives and analogs are obtained by traditional medicinal chemistry to find the best ligand or drug.

Alternatively, the structural information collected in the screening process can be used directly to suggest analogs or derivatives of the hit. This approach is illustrated when the active site is composed of one primary pocket surrounded by a variety of subsites and small pockets (Figure 1). Detailed structural information about how a compound is bound by the receptor is obtained simultaneously as a hit is detected. Such information is useful to the ordinary artisan for designing better ligands. P. Colman, Curr. Opin. in Struct.

Biology, v. 4, pp. 868-74 (1994); J. Greer et al., J. Med. Chem., v. 37, pp. 1035-54 (1994); C. Verlinde et al., Structure, v. 15, pp. 577-87 (1994). In particular, the hit identifies sites for analog synthesis which would permit access to the surrounding subsites and small pockets. This suggests the design of new compounds which better fit the active site. Furthermore, in cases where there is an existing structure-function relationship, activity enhancing substitution patterns may be directly transferred to the new lead scaffold at the 3-D structural level.

Another illustration (<u>Figure 2</u>) usually applies to a target that has two or more separate pockets that will accommodate fragments. Here, the crystalline target is screened for ligands that occupy all of the sites either in sequence or simultaneously. Because the binding event is monitored by visualizing co-crystal structures, the site of ligand binding is identified directly and there is no need for competition experiments to assure that the ligands indeed occupy different sites on the protein. Screening separately allows for

ligands which bind to distinct pockets that overlap in their binding loci. Screening for the second in the presence of the first would detect cooperative binding at a second site. Once potential leads and a structure-activity relationship have been established, linkages between each of the sites may be designed using the detailed structural information and the fragment linking approach as previously described to produce novel, much more potent ligands. S.B. Shuker et al., Science, v. 274, pp. 1531-34 (1996); C. Verlinde et al., Structure, v. 15, pp. 577-87 (1994).

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In a third application, the scaffold merging approach (not shown), the target active site is composed of two or more subsites. The crystalline protein is screened for ligand/s which bind via these subsites and the relative ligand binding orientation observed for multiple experiments. These ligands should bind by occupying one or more subsites and by overlaying the structures of multiple hits a core may be designed that will facilitate access to multiple subsites. This core would then serve as a new, novel and more potent lead compound which would also serve as the lead scaffold in the drug-design cycle.

The CrystaLEADTM linked-fragment approach experimentally implements the structure-based linked-fragment approach reported only at the computational level by Verlinde et al. in J. Comput. Aided Mol. Des., v. 6, pp. 131-47 (1992). Verlinde et al. proposed ligand fragments based on mathematical calculations. The proposed fragments were then assayed for binding activity. If the fragments actually bound, their 3-D structures were determined by X-ray crystallography and a linker designed. By contrast, CrystaLEADTM concurrently detects the binding event and provides an experimentally determined 3-D structure of the ligand-protein complex. The invention also provides for a process of determining the association constant between a target molecule and its ligand. The invention requires no special labeling of the target. Therefore, the target molecule, can encompass proteins, polypeptides, nucleic acids, nucleoproteins, or any other suitable target molecule, that is isolated from natural sources or by recombinant methods from any suitable host system as developed and practiced by the ordinary artisan.

There are several advantages to crystallographic screening. One important advantage is that the binding event is monitored directly so that the probability for false positives is reduced to near zero. The crystallographic data provide a three dimensional electron density "snap-shot" of the ligand-receptor complex showing which compound binds and how it is bound.

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The method is uniquely sensitive to structural changes in both the target and the ligand. Observing structural changes is critical in designing scaffolds which combine information from different ligand-target complex structures. One such example occurs when a protein changes structure in order to accommodate one ligand, but the structure change concurrently blocks the binding of a second ligand. Similarly, detecting structural changes is also important because if the primary scaffolds bind differently, it may not be possible to combine them into a larger scaffold.

Since the binding event is monitored directly, CrystaLEADTM does not require specially labeled samples, probes or target molecules which would be indirectly sensitive to ligand association. As long as one is able to obtain a crystal structure of the target, one can use CrystaLEADTM to screen for ligands.

If compound mixtures are suitably designed to be shape-diverse, the invention alleviates the need for de-convolution of libraries which are soaked as a mixture because the binding event is detected directly by examining the shape of the electron density at the binding site. Thus, the shape of the electron density identifies both the binding event and the compound identity directly. Alternatively, one can design the mixture to contain compounds with anomalous scattering atoms (e.g. Br, S) that can be identified by anomalous scattering techniques. Further, because CrystaLEADTM directly monitors binding, it is particularly well-suited for studying targets where no known ligand exist.

Because the electron density function calculated in CrystaLEADTM shows the "real space" of the crystal, one can focus directly on the region of interest. Thus, binding may be detected exclusively at the site of interest although the method is not limited to the active site. Binding at other sites, which complicates analysis in most binding assays, can be eliminated from consideration totally.

CrystaLEADTM also provides for a method of concurrently monitoring binding at different locations. That is, for a target with more than one pocket, screening for a second site does not require screening in the presence of the first ligand. However, screening for a second site may be completed in the presence of the first ligand in order to discover cooperative ligands.

CrystaLEAD™ is applicable for any target molecule for which a crystal structure can be obtained. According to current literature, this includes any soluble macromolecule with molecular weight between about 5000 and 200,000. However, this range expands

almost daily in response to technological advances. The method is also sensitive to a wide range of binding dissociation constants (<picomolar to molar). Using more sensitive CCD camera detectors, data may be collected in about < 4 hrs to about 4 hours with a rotating anode source. This permits the screening of thousands of compounds per detector per day. Using synchrotron sources, the number of compounds screened increase to multiple thousands per detector, and with Laue data collection methods and testing mixtures, CrystaLEADTM data can be collected in a second or less, thus permitting thousands of compounds to be tested per day per beamline. Hence, multiple detectors or a single synchrotron beamline facilitates true high-throughput screening.

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Figure 3 outlines the invention. Crystals of the target molecule are exposed to one or more compounds by soaking the crystal or by co-crystallizing the target in the presence of one or more compounds. Then crystallographic data are collected, processed and converted to electron density maps which are examined for evidence of ligand binding. One way to detect ligand binding is to compare the structure of the original crystal with the structure of the exposed crystal.

New targets may be crystallized by published conditions or by other methods well established in the art. Similarly, target structures may be available from databases such as the Protein Data Bank or could be determined by well established methodology. Advances in molecular biology and protein engineering expedite target crystallization while advances in data collection aid in rapid structure determination for targets of previously unknown structure.

Crystals that are exposed to potential ligands by soaking require an empty accessible active site. Crystals with an empty active site may be obtained by various methods, including but not limited to: crystallization in the absence of a ligand; crystallization in the presence of ligand bound at a distal site; or crystallization in the presence of a non-covalent ligand that is easily diluted or exchanged from the target once the biomolecule crystallizes. By a novel method, the inventors have obtained crystals from a biomolecule by crystallizing the biomolecule in the presence of a degradable ligand at the active site and then degrading the ligand once a crystal is formed. Alternatively, it is possible to grow the crystals in the presence of the compounds to be screened. Crystals are allowed to equilibrate in the presence of the mixture, at which point the ligands bind as a function of their concentration and binding affinity.

For the soaking method, the sensitivity of the method may be approximated by simple equilibria relationships because the concentration of protein in the crystal may be calculated and the concentration of ligand is a known quantity. For example, the concentration of a 25,000 MW protein (urokinase) in a crystal is calculated as follows: there are 4 molecules in the orthorhombic unit cell (all angles 90°) which has a volume of 55 X 53 X 82 Å³; using Avogadro's number, the concentration is 28 mM. Therefore, a mixture of compounds having a 6 mM concentration for each ligand will result in a calculated sensitivity limit of Kd < 1.5 mM (assuming a detection limit of about 80% occupancy in the crystal).

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Soaking mixtures of compounds also raises the question of multiple occupancy (more than one ligand binding to the site of interest). For cases of multiple occupancy where the ligands are bound in different pockets (see Figure 2), resolution by CrystaLEAD™ is easy because the binding at the separate sites can be distinguished individually by the electron density maps. For the scenario where different ligands compete to occupy the same site, one may use a simple competitive inhibition model to calculate the requirements for such binding. From empirical observation, it is believed that crystallography can resolve situations where the occupancy of one inhibitor is 80% and another 20%. Therefore, a ratio of binding affinity that is greater than four would result in an apparent occupancy by only the higher-affinity ligand. In the unlikely case where the ratio of binding constants of two compounds in the mixture are less than four, the resulting electron density would be a weighted average of the two separate densities and might be difficult to identify. Accordingly, it would be necessary to conduct further soaking experiments to de-convolute the mixture (e.g., looking at each compound individually in separate crystals) only where the ratio of binding affinities is less than four. This would still be worthwhile and efficient because it already determines that at least two hits are present in the mixture.

Compounds to be screened are formed into libraries. For the purposes of this discussion, libraries are large mixtures of compounds (100-10,000+) and may be general or structure-directed. A general library is random, i.e. fully diverse in size, shape and functionality. A structure-directed library is aimed at a particular functional mixture or subsite in the active site of the target molecule (e.g., a library where all compounds contain a carboxylate functionality to be directed towards a positive charge in the target active site).

In a preferred embodiment, either type of library is divided into smaller groups of shape diverse mixtures. Hence, a mixture is defined as a subset of the library which may be soaked or grown into the crystal. The mixture is determined to be shape diverse by visual inspection of the two dimensional chemical structures or computationally by programs. Shape diversity of the mixture permits a bound ligand to be identified directly from the resultant electron density map (see Figure 4). This eliminates the need for follow-up experiments to determine which compound of the mixture is a hit (bound to the target).

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If the test compounds are water soluble, typical buffers and precipitant solutions used in crystallization can be used to solubilize the mixtures and soak them into the crystal. Less water soluble compounds are dissolved individually to a final concentration of 2M in a suitable organic solvent. In one embodiment, they are dissolved in 100% DMSO and stored at 4° C, and mixed by mixing the DMSO stocks before exposure to the crystal. These mixtures would service most crystal systems where the conditions for crystal growth do not include organic reagents. The compounds would be typically soaked to a final DMSO concentration of 1-10% and allowed to equilibrate with the crystalline protein for a pre-determined amount of time (4-24 hrs). Under this scenario, each crystal is exposed to multiple compounds per soaking mixture. Some crystal growth conditions can include a high concentration of organic solvent (40-50%) which are typically alcohol derivatives. In this case, the compound libraries may be dissolved in the crystallization organic solvent which would allow a final co-solvent concentration of 40-50% for the soaking experiment. Here, the number of compounds per soaking mixture could increase.

After soaking, each crystal is exposed to a cryoprotectant such as 5-20% glycerol in the soaking mixture, mounted in a nylon loop and placed on the X-ray unit under a nitrogen cold stream (160K). The crystal studies may also be performed at room temperature or other suitable conditions as necessary for the stability of the crystals. Automated crystal mounting and changing equipment may be used to accelerate this step of the process.

30 Crystallographic data are collected and processed where each reflection (spot) on the diffraction pattern is assigned an index (h,k,l) and the intensity is measured as standard in the field. X-ray sources may be laboratory x-ray generators or high brilliance synchrotron sources that permit diffraction data collection at very high speed.

Specifically, laboratory data collection may take from 30 minutes to several hours per crystal while the time can be reduced using synchrotron sources. Data collection per crystal could be reduced to fractions of a second using Laue data collection schemes.

The diffraction data are then converted to electron density maps by methods familiar to the ordinary artisan. The electron density maps are the 3-D pictures of the ligands and/or the target biomolecules.

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For Fo-Fc maps, the calculated structure-factor amplitudes (|Fc|) which are obtained from the known crystal structure with no ligand bound are subtracted from the observed amplitudes (|Fo|). Thus, this map represents a direct subtraction of the data arising from the native protein structure from data arising from crystals soaked in the presence of a library mixture. The result is an electron density map which has positive and negative peaks. The peaks relevant to CrystaLEAD™ are the positive ones which are the direct result of ligand binding at the site of interest on the target -- that is the addition of the ligand into the target biomolecule. In Figure 3, the Fo-Fc map clearly shows a large positive peak at the active site of urokinase. The shape of the peak corresponds to the ligand 2-amino-8-hydroxyquinoline. The ligand is shown occupying the positive difference density. The other positive peaks correspond to a bound sulfate moiety (indicated by SO_4^2) and bound water molecules (indicated by H_2O). This type of map is also very sensitive to small structural changes (indicated by Δ) that, when used in conjunction with 2Fo-Fc maps, allows determination of the detailed structure of the entire ligand-protein complex. To calculate the 2Fo-Fc maps, one subtracts (|Fc|) from 2(|Fo|). Here, the map is positive and has density for all atoms of the molecule.

In <u>Figure</u> 3, inspection of the map indicates the identity and structure of the bound compound. Preferably, the maps of exposed crystals are compared with the maps of the unexposed target molecule to differentiate the positive density that may be found in the Fo-Fc map. Sometimes water molecules occupy the active site in the crystal in the absence of a bound ligand. This is easily differentiated because bound water molecules are often oriented in a geometry consistent with hydrogen bonding and because they are not connected by a network of covalent bonds. Thus, the resultant map tends to be disconnected indicating bound solvent rather than an organic compound. If the density in the Fo-Fc or 2Fo-Fc map is determined to represent an organic compound, the three-dimensional shape is compared to that of the compounds present in the library and a best-fit match is made. Alternatively, programs such as the XFIT modules of QUANTA

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(Molecular Simulations Inc., Quanta Generating and Displaying Molecules, San Diego: Molecular Simulations Inc., 1997) can automate this process.

As the ability to measure or process diffraction intensities improves, one may not need to perform the comparison on electron density maps. One may detect binding by simply comparing the diffraction patterns of the exposed crystals with the unexposed crystals. Therefore, one needs to create an electron density map only if a binding event is detected in this pre-screening process.

As shown above, CrystaLEADTM can be applied to any biomolecular target for which a crystallographic structure can be obtained. Because of its broad applicability, it is best illustrated by the examples below.

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The urokinase and VanX examples represent two scenarios for the use of CrystaLEADTM. For urokinase, the re-engineered microUK (µUK) crystals diffract very well and are of a high symmetry space group. By contrast, VanX crystals diffract more weakly and with lower symmetry. Thus, VanX requires greater data collection time. In addition, urokinase crystals have one molecule in the asymmetric unit, while VanX has six. The larger asymmetric unit requires collection of higher resolution data and makes map inspection more tedious. However, in the case of VanX, no non-substrate mimetic binders were known before those discovered by CrystaLEADTM. Therefore, CrystaLEADTM provided a novel non-peptidic lead compound to be fed into the drug-discovery cycle. For urokinase, CrystaLEADTM provided a novel primary scaffold. Applicants were able to rapidly increase the potency of the primary scaffold by using existing SAR and crystal structures to design a higher-affinity derivative with improved bioavailability over known urokinase ligands.

However, these examples illustrate the preferred embodiment of the present invention, and do not limit the claims or the specification. The ordinary artisan will readily appreciate that changes and modifications to the specified embodiments can be made without departing from the scope and spirit of the invention. Finally, all citations herein are incorporated by reference.

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EXAMPLES

Example 1: Urokinase

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5 Urokinase, a serine protease, is strongly associated with tumor cells. Urokinase activates plasminogen into plasmin which, in turn, activates the matrix metalloproteases. Plasmin and the metalloproteases degrade the extracellular matrix and promote tumor growth and metastasis. Thus, inhibitors that specifically target urokinase may serve as effective anti-cancer agents.

Human pro-urokinase consists of 411 amino acids (Figure 5). Verde et al., Proc. Nat'l Acad. Sci., v. 81(5), pp. 4727-31 (1984); Nagai et al., Gene, v. 36(1-2), pp. 183-8 (1985). When activated by proteolytic cleavage at the Lys¹⁵⁸-Ile¹⁵⁹ peptide bond, the enzyme becomes two chains connected by a single disulfide bridge (Cys¹⁴⁸-Cys²⁷⁹). The A-chain (residues 1-158) contains an EGF-like domain and a kringle domain. The B-15 chain (residues 159-411) contains the catalytic serine protease domain. Further incubation of urokinase results in an additional proteolytic cleavage at the Lys¹³⁵-Lys¹³⁶ peptide bond to form low-molecular-weight urokinase. Crystals of this enzyme form in complex with the covalent inhibitor Glu-Gly-Arg chloromethyl ketone were obtained by Spraggon et al., Structure, v. 3, pp. 681-91 (1995), and were shown to diffract to 2.5Å resolution at a high energy synchrotron source. However, the poor diffraction quality of these crystals together with the presence of a covalently bound inhibitor makes application of CrystaLEAD™ difficult.

uUK Crystal Preparation & Structure

To implement CrystaLEAD™, human urokinase was re-engineered to consist only 25 of residues 159-404 of the B-chain where Asn³⁰² was replaced with a glutamine to remove a glycosylation site and Cys²⁷⁹ was replaced with an alanine to remove the free sulfhydryl moiety. This form of urokinase (uUK) was shown to be fully active and was found to crystallize in a crystal form compatible with CrystaLEAD™. (See, also, U.S. Patent No. 30 5,112,755, issued May 12, 1992, to Heyneker et al.)

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Preparing Vector Construct pBC-LMW-UK-Ala²⁷⁹

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Mutants of human UK were cloned into a dicistronic bacterial expression vector pBCFK12. Pilot-Matias et al., <u>Gene</u>, v. 128, pp. 219-25 (1993). The following oligo nucleotides were used to generate various UK mutants by PCR:

SEQUENCE OF PCR PRIMER 1 5'-ATTAATGTCGACTAAGGAGGTGATCTAATGTTAAAATTTCAGTGTGGCCAA-3' 2 5'-ATTAATAAGCTTTCAGAGGGCCAGCCATTCTCTTCCTTGGTGTGACTCCTGATCCA-3' 5'-ATTAATTGCGCAGCCATCCCGGACTATACAGACCATCGCCCTGCCCT-3'

The initial cloning of a low molecular weight UK, hereinafter designated LMW-UK (L¹⁴⁴-L⁴¹¹) was performed using human UK cDNA as template and SEQ ID NOs: 1 and 2 as primers in a standard PCR reaction. The PCR amplified DNA was gel purified and digested with restriction enzymes *Sal*I and *Hind*III. The digested product then was ligated into a pBCFK12 vector previously cut with the same two enzymes to generate expression vector pBC-LMW-UK. The vector was transformed in DH5α cells (Life Technologies, Gaithersburg, MD), isolated and the sequence confirmed by DNA sequencing. The production of LMW-UK in bacteria was analyzed by SDS-PAGE and zymography, Granelli-Piperno et al., J. Exp. Med., v. 148, pp. 223-34 (1978), which measures plasminogen activation by UK. That LMW-UK was expressed in *E. coli*, and that it was active in the zymographic assay was demonstrated by commassie blue stained gel.

The success of the quick expression and detection of LMW-UK in *E. coli* made it possible to perform mutagenesis analysis of UK in order to determine its minimum functional structure. One mutant having a Cys²⁷⁹ to Ala²⁷⁹ replacement was made with SEQ ID Nos: 2 and 3 by PCR. The PCR product was cut with *Avi*II and *Hind* III, and used to replace a *Avi*II and *Hind* III fragment in the pBC-LMW-UK construct. The resulting pBC-LMW-UK-Ala²⁷⁹ construct was expressed in *E. coli* and the product shown to be active in zymography.

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Cloning and Expressing µUK (UK(I¹⁵⁹-K⁴⁰⁴)A²⁷⁹Q³⁰²) in Baculovirus

 μ UK (UK amino acids Ile^{159} -Lys⁴⁰⁴ that contain $Ala^{279}Gln^{302}$) was generated by PCR with the following oligonucleotide primers:

SEQ ID #	SEQUENCE OF PCR PRIMER
4	5'-ATTAATCAGCTGCTCCGGATAGAGATAGTCGGTAGACTGCTCTTTT-3'
5	5'-ATTAATCAGCTGAAAATGACTGTTGTGA-3'
6	5'-ATTAATGTCGACTAAGGAGGTGATCTAATGTTAAAATTTCAGTGTGGCCAA-3'
7	5'-ATTAATGCTAGCCTCGAGCCACCATGAGAGCCCTGCT-3'
8	5'-ATTAATGCTAGCCTCGAGTCACTTGTTGTGACTGCGGATCCA-3'
9	5'-GGTGGTGAATTCTCCCCCAATAATGCCTTTGGAGTCGCTCACGA-3'

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To mutate the only glycosylation site (Asn³⁰²) in UK, oligonucleotide primers SEQ ID NOs: 4 and 6, and SEQ ID NOs: 5 and 8 were used in two PCR reactions with pBC-LMW-UK-Ala²⁷⁹ as the template. The two PCR products were cut with restriction enzyme *Pvu* II, ligated with T4 DNA ligase, and used as template to generate LMW-UK-A²⁷⁹-Q³⁰². In the meantime, native UK leader sequence was fused directly to Ile¹⁵⁹ by PCR with SEQ ID NOs: 7 and 9 using native UK cDNA as the template.

This PCR product was used as a primer, together with SEQ ID NO: 8, in a new PCR reaction with LMW-UK-A²⁷⁹-Q³⁰² DNA as template to generate μUK cDNA. μUK was cut with *Nhe* I and ligated to a baculovirus transfer vector pJVP10z cut with the same enzyme. Vialard et al, <u>J. Virology</u>, v. 64(1); pp. 37-50, (1990). The resulting construct, pJVP10z-μUK was confirmed by standard DNA sequencing techniques.

Construct pJVP10z-µUK was transfected to Sf9 cells by the calcium phosphate precipitation method using the BaculoGold kit from PharMingen (San Diego, CA). Active µUK activity was detected in the culture medium. Single recombinant virus expressing µUK was plaque purified by standard methods, and large stock of the virus was made.

Large scale expression of μ UK was made in another line of insect cells, High-Five cells (Invitrogen, Carlsbad, CA), in suspension growing in Excel 405 serum-free medium (JRH Biosciences, LeneXa, KS) in 2 liter flasks, shaking at 80 rpm, 28°C. High-Five cells were grown to 2 X 10⁶ cells/ml, recombinant μ UK virus was added at 0.1 multiplicity of

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infection, and the culture was continued for 3 days. The culture supernatant was harvested as the starting material for purification. The activity of μ UK in the culture supernatant was measured by amidolysis of a chromogenic UK substrate S2444, which was at 6-10 mg/liter. Claeson et al., <u>Haemostasis</u>, v. 7, p. 76 (1978).

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Expressing uUK in Pichia pastoris

To express μUK in *Pichia*, an expression vector with a synthetic leader sequence was used. The *Pichia* expression vector, pHil-D8, was constructed by modifying vector pHil-D2 (Invitrogen) to include a synthetic leader sequence for secretion of a recombinant protein. The leader sequence 5'-

ATGTTCTCCAATTTTGTCCTTGGAAATTATTTTAGCTTTGGCTACTTTGCAAT CTGTCTTCGCTCAGCCAGTTATCTGCACTACCGTTGGTTCCGCTGCCGAGG GATCC-3' (SEQ ID NO: 10) encodes a PHO1 secretion signal (indicated by the single underline) operatively linked to a pro-peptide sequence (indicated in bold) for KEX2 cleavage. To construct pHil-D8, PCR was performed using pHil-S1 (Invitrogen) as template since this vector contains the sequence encoding PHO1, a forward primer (SEQ ID NO: 11) corresponding to nucleotides 509-530 of pHil-S1 and a reverse primer (SEQ ID NO: 12) having a nucleotide sequence which encodes the latter portion of the PHO1 secretion signal (nucleotides 45-66 of SEQ ID NO: 10) and the pro-peptide sequence (nucleotides 67-108 of SEQ ID NO: 10). The primer sequences (obtained from Operon Technologies, Inc. Alameda, CA) were as follows:

SEQ ID

SEQUENCE OF PCR PRIMER

- 11 5'-GAAACTTCCAAAAGTCGCCATA-3'
- 12 5'-ATTAATGAATTCCTCGAGCGGTCCGGGATCCCTCGGCAGCGGAACCAACGGTAGTGCAG
 ATAACTGGCTGAGCGAAGACAGATTGCAAAGTA-3'

Amplification was performed under standard PCR conditions. The PCR product

(approximately 500 bp) was gel-purified, cut with BlpI and EcoRI and ligated to pHil-D2

cut with the same enzymes. The DNA was transformed into E. coli HB101 cells and
positive clones identified by restriction enzyme digestion and sequence analysis. One
clone having the proper sequence was designated as pHil-D8.

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The following two oligonucleotide primers then were used to amplify μUK for cloning into pHil-D8.

SEQ ID #	SEQUENCE OF PCR PRIMER
13	5'-ATTAATGGATCCTTGGACAAGAGGATTATTGGGGGAGAATTCACCA-3'
14	5'-ATTAATCTCGAGCGGTCCGTCACTTGGTGTGACTGCGAATCCAGGGT-3'

The PCR product was obtained with SEQ ID NOs: 13 and 14 using pJVP10z-μUK as the template. The amplified product was cut with BamHI and XhoI and ligated to pHil-D8 cut with the same two enzymes. The resulting plasmid, pHilD8-μUK, was confirmed by DNA sequencing, and used to transform a *Pichia* strain GS115 (Invitrogen) according to the supplier's instructions. Transformed *Pichia* colonies were screened for μUK expression by growing in BMGY medium and expressing in BMMY medium as detailed by the supplier (Invitrogen). The μUK activity was measured with chromogenic substrate S2444. The μUK expression level in *Pichia* was higher than that seen in baculovirus-High Five cells, ranging from 30-60 mg/L.

15 Purifying μUK

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The culture supernant of either High Five cells or *Pichia* were pooled into a 20 liter container. Protease inhibitors, iodoacetamide, benzamidine and EDTA were added to a final concentration of about 10 mM, 5 mM and 1 mM, respectively. The supernatant was then diluted 5-fold by adding 5 mM Hepes buffer pH7.5 and put through 1.2 μ and 0.2 μ filter membranes. The μUK was captured onto Sartorius membrane adsorber S100 (Sartorius, Edgewood, NY) by passing through the membrane at a flow rate of 50~100 ml/min. After extensive washing with 10 mM Hepes buffer, pH7.5, 10 mM iodoacetamide, 5 mM benzamidine, 1 mM EDTA, μUK was eluted from S100 membrane with a NaCl gradient (20 mM to 500 mM, 200 ml) in 10 mM Hepes buffer, pH7.5, 10 mM iodoacetamide, 5 mM benzamidine, 1 mM EDTA. The eluate (~100ml) was diluted 10 times in 10 mM Hepes buffer containing inhibitors, and loaded to a S20 column (BioRad, Hercules, CA). μUK was eluted with a 20x column volume NaCl gradient (20 mM to 500 mM). No inhibitors were used in the elution buffers. The eluate was then diluted 5-fold with 10 mM Hepes buffer, pH7.5, and loaded to a heparin-agarose (Sigma) column.

μUK was eluted with a NaCl gradient from 10 mM to 250 mM. The heparin column eluate of μUK (~50 ml) was applied to a benzamidine-agarose (Sigma, St. Louis, MO) column (40 ml) equilibrated with 10 mM Hepes buffer, pH7.5, 200 mM NaCl. The column was then washed with the equilibration buffer and eluted with 50 mM NaOAc, pH 4.5, 500 mM NaCl. The μUK eluate (~30 ml) was concentrated to 4 ml by ultrafiltration and applied to a Sephadex G-75 column (2.5x48 cm; Pharmacia Biotech, Uppsala, Sweden) equilibrated with 20 mM NaOAc, pH4.5, 100 mM NaCl. The single major peak containing μUK was collected and lyophilized as the final product. The purified material appeared on SDS-PAGE as a single major band.

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High-quality µUK crystals facilitated determination of its apo-three-dimensional structure by X-ray crystallography to 1.0Å resolution. Crystals were obtained by the hanging drop vapor diffusion method. Typical well solutions consisted of 0.15M Li₂SO₄, 20% polyethylene glycol MW 4000 and succinate buffer pH 4.8-6.0. On the cover slip, 2 · µl of well solution were mixed with 2 µl of protein solution and the slip sealed over the well. Crystallization occurred at approximately 18-24°C within 24 hrs. The protein solution contained 6 mg/ml (0.214mM) μUK in 10 mM citrate pH 4.0, 3 mM ε-amino caproic acid p-carbethoxyphenyl ester chloride (inhibitor) with 1% DMSO co-solvent. The inhibitor utilized in the co-crystallization is believed to acylate the active site serine 195 and is subsequently deacylated enzymatically, because, the 3-D X-ray structure of crystals grown in the presence of this compound show no inhibitor remaining in the enzyme active site. Menegatti et al., J. Enzyme Inhibition, v. 2, pp. 249-59 (1989). The only density present is that due to bound solvent molecules. Because $\mu U K$ will not crystallize in the absence of the inhibitor, the meta-stable inhibitor:UK complex is believed to be the crystallization entity. Importantly, the resultant μUK crystals are composed of enzyme with an empty active site which is the ideal case for implementation of CrystaLEAD™.

Crystals obtained under these conditions belong to the space group P2₁2₁2₁ with unit cell dimensions of a=55.16Å b=53.00Å c=82.30Å and $\alpha=\beta=\gamma=90^{\circ}$. They diffract to beyond 1.5Å on a rotating anode source. Further, a 1.0Å resolution native data set has been collected at the Cornell High Energy Synchrotron Source in Ithaca, New York. The crystal structure was determined by the molecular replacement method using the AMORE program, Navaza, <u>J. Acta Cryst.</u>, A50:157-163 (1994), with the low-resolution urokinase structure as the search probe, Spraggon et al., <u>Structure</u>, v. 3, pp. 681-691 (1995); PDB

entry 1LMW. The structure was refined using the XPLOR program package, A. Brunger, X-PLOR (version 2.1) Manual, Yale University, New Haven CT (1990).

Screening for Weak Bases

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The µUK was screened against a structure-directed library in order to find a novel primary scaffold which would have favorable pharmacokinetic properties. Since the urokinase active site is composed of one primary pocket that contains a free carboxylate moiety in the form of an aspartic acid (Asp¹⁸⁹), most well-known scaffolds are strongly basic and contain amidine or guanidine moieties. The basic group has been found to hydrogen bond salt-link with Asp¹⁸⁹. This can be a problem pharmacologically since strong bases are known to decrease oral bioavailability. Accordingly, a weakly basic library containing compounds that were not previously known to be urokinase binders was selected.

A weak base library containing 61 compounds with pKa between about 1 and 9 was located in the available chemicals directory (ACD). The library was broken down into 9 mixtures of about 6 to 7 shape-diverse compounds, as determined by visual inspection of the two dimensional chemical structure. The compound mixtures were screened by the method described above. Specifically, each compound was dissolved in 100% DMSO to a final concentration of about 2M (or saturation for the less soluble). Equal volumes of each of the 6 or 7 compounds comprising the mixture were mixed to a final individual compound concentration of 0.33M. Single μUK crystals were placed in 50ml of 27% PEG4000, 15.6mM succinate pH 5.4, 0.17M Li₂SO₄ and 0.5-0.8ml of the compound mixture added to give 1 to 1.6% DMSO and 3.3 to 5.2mM final individual compound concentration. Under these conditions the sensitivity of the experiment is expected to detect binders with Kd<1.0mM. Crystals were allowed to equilibrate for about 8-24hrs.

Data were collected on a Rigaku RTP 300 RC rotating anode source with a RAXISII or MAR image plate detector. Typical data consisted of 45-50 2° oscillations with 2-5min exposures. Typical data were 70-90% complete at 2.0-3.0Å resolution with merging R-factors of 13-26%. Hence, the data quality ranged from fair to poor due to the rapid data collection protocol. However, this quality of data was shown to be adequate for the detection of binders primarily due to the high quality of the starting model which had

been refined to 1.5Å resolution (R=20.7% R_{free}=25.3%). Data were processed by the DENZO program package, Otwinowski et al., <u>Methods in Enzymology</u>, 276 (1996), and the electron density maps calculated by the XPLOR package.

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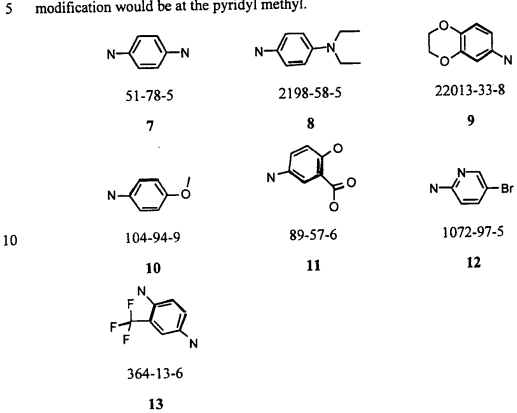
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Electron density maps were inspected on a Silicon Graphics INDIGO2 workstation using the QUANTA 97 program package (Molecular Simulations Inc., Quanta Generating and Displaying Molecules, San Diego: Molecular Simulations Inc., 1997). The shape of the density at the active site was visually identified as resulting from one (or more) of the compounds in the mixture indicating a positive hit or from ordered water molecules indicating the absence of binding. For experiments which resulted in a positive hit, the appropriate compound was visually moved into the electron density. The electron density maps were also checked for any changes in the protein structure and if observed, the appropriate modifications were made. Hence, after the map inspection/compound fitting step, the three-dimensional structure of the compound:protein complex was known. The urokinase example utilized visual movement of the compound into the density because the screening was still on a small scale. When expanded to larger scale compound screening, commercial programs such as the XFIT module of QUANTA will facilitate automatic fitting of the compound to the density.

Figure 6 shows an example of a positive hit. The compounds screened are numbered 1 through 6 and the Fo-Fc electron density map at the active site is shown at Figure 6A. The shape of the density identified the binder as compound 5. Figure 6B

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shows the detailed binding mode of the compound in the primary specificity pocket as obtained directly by interpretation of the CrystaLEADTM electron density map. The amino nitrogen hydrogen bonds with the Asp¹⁸⁹ carboxyl and the pyrimidyl nitrogen hydrogen bonds with a backbone carbonyl (Gly²¹⁸). The structure also shows that the ideal site for modification would be at the pyridyl methyl.



Another mixture of compounds (compounds 7 through 13) did not produce any hits. The resulting electron density map after soaking this group did not correspond to that of any of the tested compounds in this mixture. Instead, they correspond to bound solvent molecules. See <u>Figure</u> 6C.

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Figure 7 shows another example of a positive hit. Of the seven compounds screened (14-20), the Fo-Fc map shown in Figure 7A indicates that compound 19 is bound. The binding mode depicted in Figure 7B shows that the 2-amino is hydrogen bonding with the Asp189 side chain and that the 8-hydroxyl is an ideal site for substitution in order to access the adjacent hydrophobic sub-pocket (denoted as S1β in Figure 7B). Figure 8 represents another hit where compound 22, 5-aminoindole, (Figure 8A) was found to bind to urokinase with the amino group hydrogen bonding with Asp189 (Figure 8B). Compounds screened were compounds 21-27.

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Figure 9 shows an example where two compounds from the same mixture (compounds 28-34) were found to bind without multiple occupancy problems. In the 5 initial experiment where the crystal was soaked in the presence of the entire compound mixture, compound 28 was found to bind (Figure 9A). In addition, when the weaker binding compound 31 was soaked individually (based upon previous structure activity relationships established through CrystaLEAD™) it was also found to bind (Figure 9B). In a more typical application of the method, a library would be re-soaked in the absence of 10 the tighter binder in order to detect weaker binders in the mixture, if desired.

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Table 1 summarizes the inhibition constants for each of the CrystaLEAD[™] hits as determined by pyroGlu-Gly-Arg-pNA/HCl (S-2444, Chromogenix) chromogenic activity. Assays were completed at both pH 6.5 (0.1 M NaPO₄) and 7.4 (50mM Tris). Other conditions of the assay were 150mM NaCl, 0.5% Pluronic F-68 detergent, 200mM S-2444, with a final DMSO concentration of 2.5%. The Km of the substrate was determined to be 55μM.

Table 1: Inhibition constants and pKa for hits detected by CrystaLEAD™

Compound (CAS #)	Ki	Ki	pKa
	(pH 6.5)	(pH 7.4)	
5 (42753-71-9)	>>500µM	>>500µM	6.0*
19 (70125-16-5)	56μΜ	137μΜ	7.3
22 (65795-92-8)	200μΜ	>500µM	6.0*
28 (580-22-3)	$71 \mu M$	136μΜ	7.3
31 (1603-41-4)	>>500µM	>>500µM	7.0*

10 • indicates estimated pKa

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Based upon the activity and structural information, compound 19 was chosen as the lead compound. Crystallographic information indicated that substitution at the 8-position should allow access to the adjacent hydrophobic pocket (S1β) pocket and thereby result in an increase in potency. Based upon crystallographic and binding information from an amidine-based series, compound 35 was synthesized (the 8-aminopyrimidinyl analog of compound 19). This modification resulted in about a 200 fold increase in binding potency at pH 6.5 (Ki pH7.4=2.5μM; Ki pH6.5=0.32μM). The experiment indicates that CrystaLEADTM can provide both a lead scaffold and the detailed structural information necessary to elaborate that scaffold through structure-based drug design into a more potent compound.

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In <u>Figure</u> 10, an overlay of the crystal compound 35:urokinase and the parent compound 19 are shown. The overlay shows that the aminopyridine ring is bound in the hydrophobic sub-pocket (S1β) pocket as predicted and that this substitution results in movement of the quinoline ring towards this site.

Compound 35, the 8-aminopyrimidinyl-2-aminoquinoline, was also tested for oral bioavailability. Compound 35 was determined to be 30-40% orally bioavailable in the rat when administered at a 10mg/kg dose. Hence, successful implementation of CrystaLEADTM resulted in a novel lead scaffold which through one cycle of structure-based drug design produced a compound having a 200-fold increase in potency, and was found to be orally bioavailable.

Example 2: VanX

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Vancomycin is the drug of choice for infections caused by streptococcal or staphylococcal bacterial strains that are resistant to β-lactam antibiotics. However, strains of vancomycin resistant bacteria have now been found for this drug of last recourse. Some investigators have associated VanX, a metalloproteinase, with vancomycin resistance. VanX is part of a cascade that results in replacement of the terminal D-Ala-D-Ala moiety of the bacterial peptidoglycan chain (the binding site for vancomycin) with a D-Ala-D-lactate. This results in a 1000-fold decrease in vancomycin binding. The only known inhibitors of VanX are peptides or peptide derivatives, such as phosphonate or phosphinate analogs of the D-Ala-D-Ala substrate. As such, they are not suitable drugs because they are metabolized and/or degraded in vivo. Initial attempts to find suitable drugs by normal screening methods did not uncover a suitable ligand. Subsequently, Applicants turned to CrystaLEADTM to find a non-peptide lead compound for drug development towards a treatment for these resistant strains.

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VanX Preparation

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E. coli W3110 containing plasmid pGW1, in which the vanX gene is under control of the IPTG-inducible tac promoter, was grown at 37 °C in LB medium containing ampicillin (100 μ g/ml) to an absorbance of about 1.3-1.5 at 595 nm. Then IPTG was then added to a final concentration of 0.8 mM, and the cells were grown for an additional 1.5 hours.

Cells were harvested by centrifugation at 6000 rpm for 10 min. Then, the pellet was resuspended in ice cold 20 mM Tris-HCl (pH 8.0) containing 0.01% NaN₃, 1 mM MgCl₂, 1mM PMSF, 1 mM DTT (Buffer A) and 25 units/ml of benzonase (Nicomed Pharma, Copenhagen, Denmark). The cells were lysed by the addition of 0.1 micron zirconia ceramic beads to the lysate mixture (1:1 v:v) with a 1-3 minute run in a Bead Beater (Biospec), an ultrasound bead mill. The Bead Beater was run with an ice-packed reservoir to maintain a chilled lysate. Then, the lysate was decanted away from the settled glass beads. The beads were then rinsed with 1-2 volumes of lysis buffer, and the washes were then pooled with the original lysate. The lysate was centrifuged at 25000g for 30 minutes to settle cell debris. The supernatant was dialyzed overnight at 4 °C in 50mM Tris-HCl, pH 7.6, 1 mm EDTA, and 1mM DTT (Buffer B).

Thereafter, the dialyzed lysate was loaded onto a Q-sepharose fast flow column, pre-equilibrated in Buffer A at a rate of four millimeters per minute. The column was exhaustively washed with the Buffer A followed by a linear gradient of Buffer B to Buffer B+0.5 M NaCl. The active VanX fractions from this step were pooled, concentrated and then applied to a Superose-75 column in Buffer B. VanX fractions from the Superose column run were then applied to a Source-Q column in Buffer A at a flow rate of 2 ml/min. The column was washed with starting buffer for several column volumes. Then the VanX protein was eluted off with a shallow gradient of Buffer A to Buffer A+25mM NaCl. The active VanX fractions from this final step were concentrated to a final concentration of approximately 15 mg/ml in Buffer A with Amicon filters. Unless otherwise specified, the foregoing procedure was run at 4 °C. As purified, the VanX protein was approximately 95% pure and readily crystallized.

VanX Crystal Structure

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The crystal structure of VanX was determined at 2.2 Å resolution by multiple isomorphous replacement. Bussiere *et al.*, Molecular Cell, Vol. 2, pp 75-84 (1998). The recombinant protein obtained above was crystallized in the space group P2₁ by the sitting drop vapor diffusion method. Typical crystals had unit cell dimensions of a=83.4Å, b=45.5Å, c=171.4Å, α=γ=90°, β=104° with six molecules in the asymmetric unit. Typical well solutions consist of 0.1M Mes pH 6.4, 0.24M ammonium sulfate, and 20% PMME 5000. On the sitting drop microbridge (Hampton USA), 2ml of protein are mixed with 2ml of well solution and the chamber sealed with a cover slip. Crystallization occurs at 18°C, and the crystals grow to full size in about 2-3 days. The protein solution is composed of 12-15mg/ml (0.5-0.6mM) VanX in 10mM Tris, 15mM DTT, pH 7.2. The 3-D structure for crystals grown under these conditions show an empty active site making this a system highly suitable for application of CrystaLEADTM.

The VanX active site has an extended pocket capable of accommodating the D-Ala-D-Ala substrate. The pocket also contains a catalytic zinc. Thus, for this case, VanX was initially screened against zinc directed libraries in order to find multiple binding scaffolds which could be merged into a single lead compound. Three libraries utilizing amino-acid, thiol, hydroxamic acid or carboxylate moieties directed towards zinc were screened.

Screening

The amino acid library consisted of 102 compounds of optically pure commercially available natural and non-naturally occurring amino acids. The library was divided into 12 mixtures of 8-10 shape-diverse compounds and screened by the method described above. Specifically, each compound was dissolved in 100% DMSO to a final concentration of 2M (or saturation for the less soluble). Equal volumes of each compound of each mixture were mixed to a final individual compound concentration of 0.33M. Single VanX crystals were placed in 50ml of 0.1M Mes pH 6.4, 0.24M ammonium sulfate, 20% PMME 5000 and 0.5-0.8ml of the compound mixture added to give 1 to 1.6% DMSO and 3.3 to 5.2mM final individual compound concentration. Crystals were

allowed to equilibrate for 3-4 hrs. The thiol, hydroxamic and carboxylate libraries were prepared and screened in a similar manner.

Data were collected on a Rigaku RTP 300 RC rotating anode source with a RAXISII, MAR image plate, or MAR CCD detector. For the image plate systems, typical data consisted of 90 1.25° oscillations with 15min exposures while for the CCD 100 1.0° oscillations were exposed for two minutes. Typical usable data were >90% complete at 2.6-2.8Å resolution with merging R-factors of 10-20%. This was required to adequately visualize and identify inhibitors in the Fo-Fc or 2Fo-Fc maps. For these maps, the starting model had been refined to 2.1Å resolution (R=25% R_{free}=28%). Data were processed by the DENZO program package and the electron density maps calculated by the XPLOR package. In the presence of some compounds of the carboxylate library, the space group was shown to shift from P21 to C2 (a=170.6Å, b=47.5Å, c=83.6Å, α = γ =90°, β =104°). For this form, the asymmetric unit contained a trimer thereby reducing the number of degrees of freedom so that lower resolution data (3.0Å) were adequate for visualization of binding.

Electron density maps were inspected on a Silicon Graphics INDIGO2 workstation using QUANTA 97. The shape of the density at the active site was visually identified by the shape of one or more of the compounds in the mixture to indicate a positive hit or by ordered water molecules indicating the absence of binding. For experiments which resulted in a positive hit, the appropriate compound was visually moved into the electron density. The electron density maps were also checked for any changes in the protein structure, and if observed, corresponding modifications were made in the structure. Hence, after the map inspection/compound-fitting step, the detailed 3-D structure of the compound:protein complex was known.

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Currently 6 hits have been detected in the VanX screens (compounds 36-41).

Figure 11 shows the binding mode of representative hits. In all cases, the electron density shape identified the binding compound. Figure 11A shows compound 39 bound with the carboxylate coordinating to the active site zinc. Figure 11B, shows compound 36 bound with the carboxylate pointing towards the active site zinc. In Figure 11C, compound 37 was also found to bind through the carboxylate. The binding of compound 39 and compound 41 (not shown) suggests that the active site zinc prefers coordination of a carboxylate over a free thiol. This led to screening of a carboxylate library where additional hits were found. In all cases, the compounds were screened in mixtures of 7-10 and the hit directly identified by the shape of the electron density map. These hits are fed directly into the structure-based drug design cycle in a manner similar to that described for the urokinase example.

Example 3: Screening with Mixtures of 100 Compounds

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In order to increase the number of compounds that may be screened per unit time by the CrystaLEADTM method, a preferred embodiment of the method would be to screen mixtures of 100 compounds rather than mixtures of 10. The advantage of this method is a higher compound throughput with a concurrent lowering of the sensitivity of the hit detection. In addition, since only the most potent compound in a mixture will bind, weaker hits may be missed. When a general library, for example, one which is fully diverse in size, shape and functionality, is screened by CrystaLEADTM, the hit-rate is expected to be low. Therefore, a more coarse screen is warranted. In addition, since the hits from this screen would be the more potent binders, they could serve as starting

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scaffolds for structure-based drug design. Since the compound mixture will be composed of 100 compounds, the mixture should be carefully designed in order to ensure that all members would be diversely shaped enough to eliminate the need for deconvolution. Hence, upon hit detection, some deconvolution may be necessary to identify the hit.

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To test this particular method, a compound known to bind to μUK was added to a group of 100 compounds. This known binder, compound 19, was originally discovered by the CrystaLEADTM method and shown to bind to μUK with a Ki of 56μM at pH 6.5 and 137μM at pH 7.4. The 100 compound mixture was constructed by mixing 10 mixtures of 10 compounds. Specifically, each dry mixture of 10 was dissolved in 100% DMSO to a final concentration of about 80-240mM(or saturation for the less soluble). Equal volumes of each of the mixtures of 10 compounds were mixed to a final individual compound concentration of 8.0-24.0mM and the mixture spiked with a 100% DMSO stock of compound 19 such that the final concentration was 18.0mM. Single μUK crystals were placed in 50μl of 27% PEG4000, 15.6mM succinate pH 5.4, 0.17M Li₂SO₄ and 0.5μLof the compound mixture added to give 1% DMSO. The final concentration of each compound in the soak experiment ranged from 80-240μM, the concentration of compound 19 was 180μM. Under these conditions the sensitivity of the experiment is expected to detect binders with Kd<20-60μM. Crystals were allowed to equilibrate for 4 hours and 15 minutes.

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Data were collected at the Argonne National Labs advanced photon source synchrotron ID beamline IMCA equipped with a MarCCD camera. Data consisted of 100 1° oscillations with 7 sec exposures. Data were 87.4% complete at 1.6Å resolution with an overall merging R-factor of 5.4%. Data were processed by the DENZO program package, Otwinowski et al., Methods in Enzymology, 276 (1996), and the electron density map calculated by the XPLOR package.

The electron density map was inspected on a Silicon Graphics INDIGO2 workstation using the QUANTA 97 program package (Molecular Simulations Inc., Quanta

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Generating and Displaying Molecules, San Diego: Molecular Simulations Inc., 1997). The shape of the density at the active site was visually identified as resulting from one of the compounds in the mixture indicating a positive hit which was identified as compound 19, and is illustrated in Figure 12.

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This method is preferable for discovering lead compounds. Lead compounds would typically have the characteristics of being tighter binders (for example, within the sensitivity range of the method). This method also allows screening of a 10,000 compound non-directed library on the timeframe of 1-2 weeks. This method would be used in conjunction with the other methods of screening 10-20 compounds at a time where weaker binders would be identified. These binders would be less likely to serve as lead compounds, but could be attached to a lead scaffold in order to increase the potency.

15 Example 4: CrystaLEAD™ screening of ErmC'

ErmC' is an rRNA methyltransferase that transfers a methyl group from S-Adenosyl-L-methionine to N6 of adenine within the peptidyltransferase loop of 23S rRNA. This methylation confers antibiotic resistance against a number of macrolide antibiotics such as the widely prescribed erythromycin. Inhibition of ErmC' would be expected to reverse resistance. In order to design a specific and potent ErmC' inhibitor, the cofactor or S-Adenosyl-L-methionine binding site has been targeted. S-Adenosyl-L-methionine is illustrated below as compound 42:

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The crystal structure of ErmC' shows that the S-Adenosyl-L-methionine site is composed of two primary pockets which accommodate the adenine ring and the methionine. In addition, there is a third pocket which may accommodate the rRNA adenine that undergoes methylation. In order establish an SAR at this site, a library of adenosine analogues substituted at N6 and/or 5' hydroxyl was generated. The sites of variation in the library is represented below as compound 43.

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ErmC expression and purification

The expression vector pTERM31 was constructed by polymerase chain reaction

(PCR) amplification of the ermC' gene and the upstream kdsB cistron from pERM-1.

Subcloning the PCR product into pET24+ (Novagen, Madison, WI) was performed using the *Bam*HI and *Hin*dIII sites included in the "tailed" PCR primers. This new construct allowed the expression of ErmC' by translational coupling to kdsB, under the control of the T7*lac* promoter. pTERM31 plasmid was transformed into *E. coli* strain

BL219(DE3)/pLysS (Novagen) and the resulting strain was used for production of ErmC'. Transformed cells were grown at 27.5°C in a New Brunswick Scientific (Edison, NJ)

Micros fermentor containing 101 of Superbroth (BIO 101, La Jolla, CA), supplemented with kanamycin, chloramphenicol, and glucose. When the culture optical density reached 1.10, ErmC' expression was induced by the addition of 1mM isopropyl β-d-thiogalactopyranoside (IPTG). Cells were harvested 400 minutes post-induction.

Frozen cell paste (200-250g) was thawed at room temperature and resuspended into 5-10 volumes of cold lysis buffer (50mM Tris, 5mM 1,4-dithiothreitol (DTT), 1mM phenylmethylsulfonate fluoride (PMSF), 2mM ethylene-diaminetetraacetic acid (EDTA), 0.2% Triton X-100, ph 7.8). The cells were lysed with a French press and cell debris removed by centrifugation. The supernatant was dialysed overnight against 20 l Tris-DTT-glycerol-magnesium (TDGM) buffer, pH 7.8 (50mM Tris, 5mM DTT, 10% glycerol, 10mM MgCl2). The dialysate was then applied to a Sepharose Fast Flow column (Pharmacia) that had been pre-equilibrated in TDGM buffer. Fractions were assayed for methyltransferase activity and those containing ErmC' were pooled, applied to a TSK SP-5PW column (TosoHaas, Montgomeryville, PA), and eluted with an

NaCl gradient. The purified protein was then concentrated on a YM-10 (Amicon) membrane.

ErmC Crystal Structure

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Crystals of ErmC' were grown by the hanging drop vapor diffusion method. Drops containing 5-8 mgs/ml ErmC' in 25mM Tris/Cl, 100mM NaCl, 2mM DTT, 10% (v/v) glycerol, pH 7.5 were equilibrated against a reservoir containing 100mM Tris, 500mM NH4(SO)4, 15% PEG 8000, pH 7.8. Crystals appeared within one day and grew to their full size within one week. Crystals belonged to the space group P43212. The

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structure of ErmC' in this space group was determined by molecular replacement to 2.2 angstrom resolution using the crystal structure of ErmC' in the space group P6 (Bussiere et al., Biochemistry Vol. 37, pp 7103-7112). The 3-D structure for crystals grown under these conditions show an empty active site making this a system highly suitable for application of CrystaLEADTM.

Screening

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The adenosine library consisted of 59 compounds. The library was divided into 7 mixtures of 8-9 shape-diverse compounds and screened by the CrystaLEADTM method. Specifically, each compound was dissolved in 100% DMSO to a final concentration of 1M (or saturation for the less soluble). Equal volumes of each compound were mixed to assemble the mixture of 10. Single ErmC crystals were placed in 50μl of 20% PEG 8000, 0.3M ammonium sulfate, 10% glycerol, pH 7.7 and 0.5-0.8μl of the compound mixture added to give 1 to 1.6% DMSO and 3.3 to 5.2μM final individual compound concentration. Crystals were allowed to equilibrate for 3-4 hrs.

Data were collected on a Rigaku RTP 300 RC rotating anode source with a RAXISII, MAR image plate, or MAR CCD detector. For the image plate systems, typical data consisted of 15-20 2° oscillations with 20-30min exposures while for the CCD 15-20 2.0° oscillations were exposed for 8-15 minutes. Typical usable data were 80-90% complete at 3.4-3.6Å resolution with merging R-factors of 7-16%. This was required to adequately visualize and identify inhibitors in the Fo-Fc or 2Fo-Fc maps. For these maps, the starting model had been refined to 2.2Å resolution (R=22% Rfree=25%). Data were processed by the DENZO program package and the electron density maps calculated by the XPLOR package.

Electron density maps were inspected on a Silicon Graphics INDIGO2 workstation using QUANTA 97. The shape of the density at the active site was visually identified by the shape of one or more of the compounds in the mixture to indicate a positive hit or by ordered water molecules indicating the absence of binding. For experiments which resulted in a positive hit, the appropriate compound was visually moved into the electron density. The electron density maps were also checked for any changes in the protein structure, and if observed, corresponding modifications were made in the structure.

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Hence, after the map inspection/compound-fitting step, the detailed 3-D structure of the compound:protein complex was known.

Two hits were detected in the ErmC' adenosine analogue screen (compounds 44 and 45). Figures 13 and 14 show the crystal structure of the complexes of compounds 44 and 45 with ErmC'.

In all cases, the electron density shape identified the binding compound. The
hydrophobic substitution was found to bind along a partially exposed hydrophobic surface
suggesting a preferred interaction which may have contributed to the binding of these
compounds, allowing them to be pulled out as hits. No hits containing a substitution at the
5'OH position were detected. A follow-up compound to compounds 44 and 45 contained
an optimized indane substituent at this hydrophobic site.

WE CLAIM:

- 1. A process for identifying a ligand to a target biomolecule comprising.
 - a) obtaining a target biomolecule crystal;
 - b) exposing the target biomolecule crystal to one or more test samples; and
 - c) obtaining an X-ray crystal diffraction pattern to determine whether a ligand/receptor complex is formed.
- 2. The process according to Claim 1 further comprising the steps of obtaining an X-ray crystal diffraction pattern of the target biomolecule crystal prior to exposure to the test samples and comparing the X-ray diffraction pattern of the target molecule before and after the exposure.
- 3. The process according to Claim 1 further comprising the step of transforming diffraction pattern into an electron density map.
 - 4. The process according to Claim 3 further comprising the step of converting electron density map into a structure.
- 20 5. The process according to Claim 1, wherein the target biomolecule is exposed to a test sample by soaking the target biomolecule crystal in a solution that contains the test sample.
- 6. The process according to Claim 1, wherein the target biomolecule is exposed to the test samples by soaking the target biomolecule crystal in a solution containing a mixture of test samples.

- 7. The process according to Claim 1, wherein the target biomolecule is exposed to the test sample by co-crystallizing the target biomolecule crystal with a test sample.
- 8. The process according to Claim 1, wherein the target biomolecule is exposed to the test samples by co-crystallizing the target biomolecule crystal with a mixture of test samples.
 - 9. The process according to Claim 6, wherein the mixture of test samples are diversely shaped.
 - 10. The process according to Claim 8, wherein the mixture of test samples are diversely shaped.
 - 11. The process according to Claim 1 wherein the ligand is a biologically-active moiety.
 - 12. The process according to Claim 1, wherein the target is a polypeptide.
 - 13. The process according to Claim 1, wherein the target is a re-engineered polypeptide.
- 20 14. A biologically-active moiety identified by the process according to Claim 11.
 - 15. The process according to Claim 1 wherein said ligand is a lead compound.
 - 16. A process to design a ligand for a target biomolecule comprising,
- 25 a) obtaining a target biomolecule crystal;

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b) identifying at least two ligands to the target biomolecule by X-ray crystallographic screening;

- c) determining the spatial orientation of the ligands when they are bound to the target biomolecule; and
- d) linking the ligands together according to the spatial orientation to form the ligand.

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- 17. The process according to Claim 16 wherein the spatial orientation of the bound ligands is determined by forming a multi-ligand/target molecule complex and generating an X-ray crystal structure of the multi-ligand/target molecule complex.
- 10 18. The process according to Claim 16 wherein one ligand is bound to the target molecule before another ligand is bound to the target molecule.
 - 19. The process according to Claim 16 wherein the ligand is a biologically-active moiety.

- 20. The process according to Claim 16, wherein the target is a polypeptide.
- 21. The process according to Claim 16, wherein the target is a re-engineered polypeptide.
- 20 22. A biologically-active moiety designed by the process according to Claim 19.
 - 23. The process according to Claim 16 wherein said ligand is a lead compound.
 - 24. A process to design a ligand for a target biomolecule comprising,
- 25 a) obtaining a target biomolecule crystal;
 - b) identifying a ligand to the target biomolecule by X-ray crystallographic screening;

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- c) making derivatives of the ligand.
- 25. The process according to Claim 24 wherein said ligand is a lead compound.
- 5 26. The process according to Claim 24 wherein the ligand is a biologically-active compound.
 - 27. The process according to Claim 24, wherein the target is a polypeptide.
- 10 28. The process according to Claim 24, wherein the target is a re-engineered polypeptide.
 - 29. A lead compound identified by the process of Claim 25.
 - 30. A biologically-active compound designed by the process according to Claim 25.
 - 31. A biologically-active compound designed by the process according to Claim 26.
 - 32. A process to form a crystal having an easily accessible active site from a biomolecule comprising,
- 20 a) co-crystallizing the biomolecule with a degradable ligand; and
 - b) degrading the ligand once the crystal is formed.
 - 33. The process according to Claim 32 wherein the biomolecule active site degrades the ligand.

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- 34. The process according to Claim 32 further comprising adding degradation agents to degrade the ligand.
- 35. The process according to Claim 32 wherein said ligand spontaneously degrades.

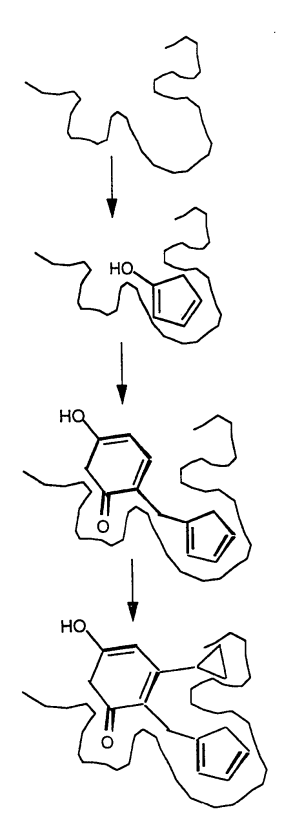


Figure 1

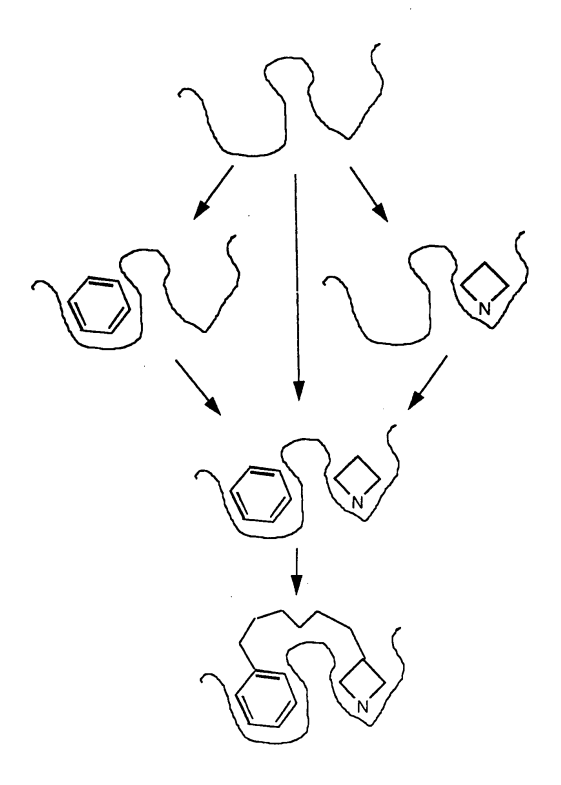


Figure 2

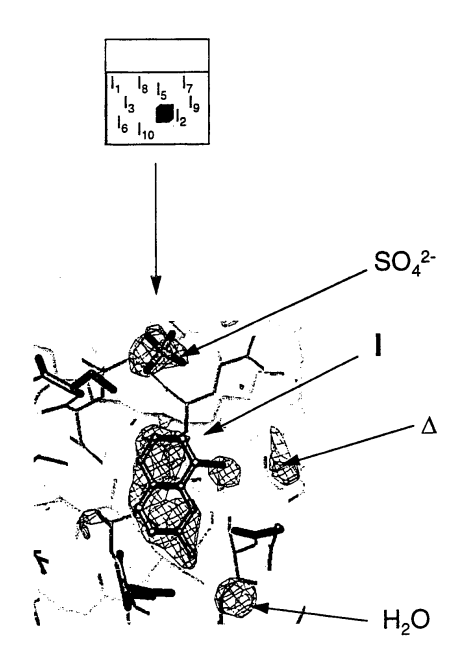


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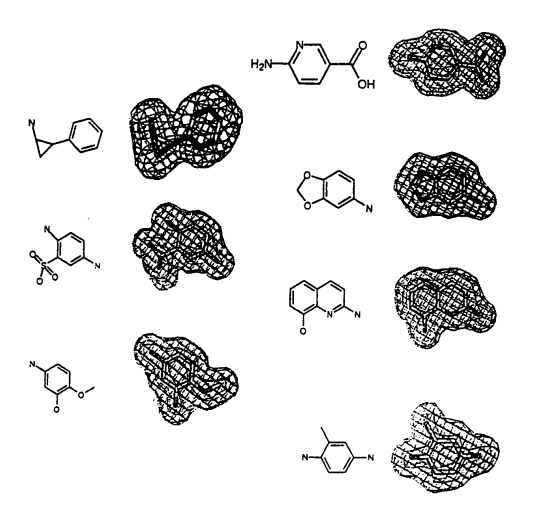
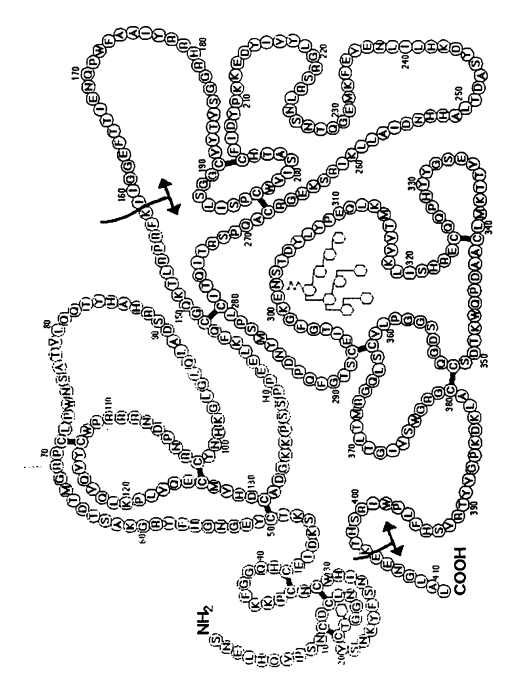
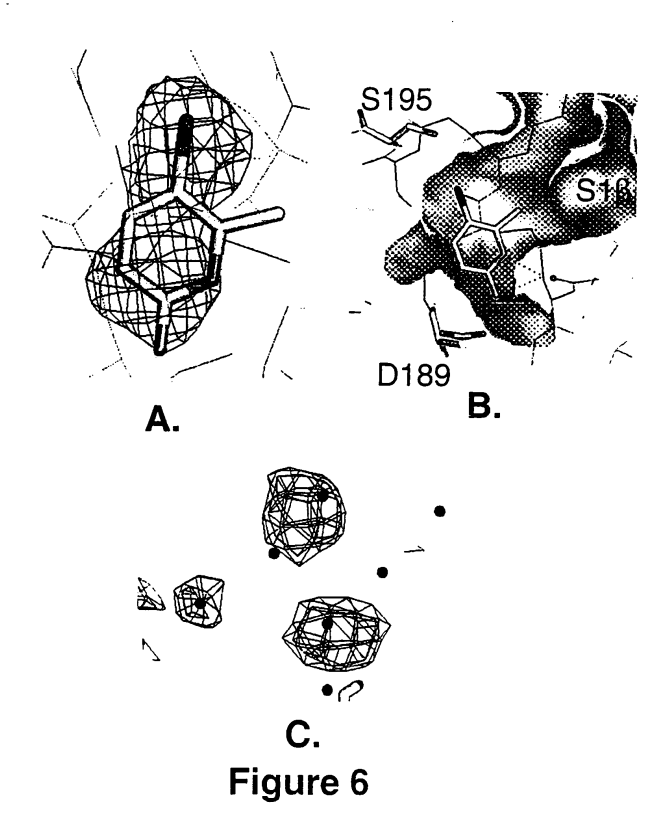
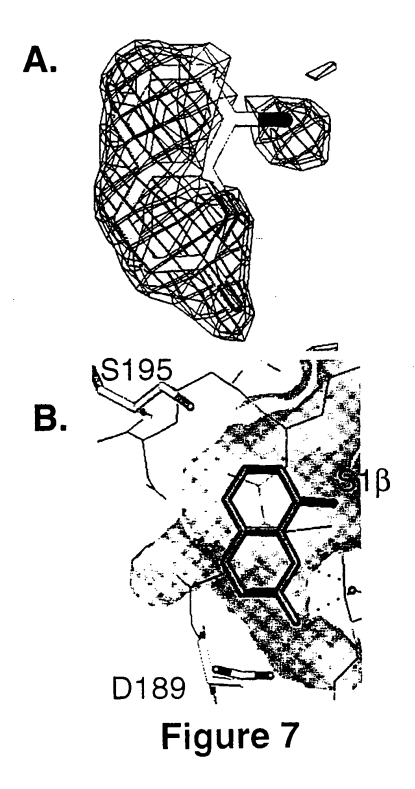


Figure 4





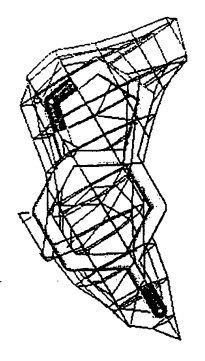




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B.

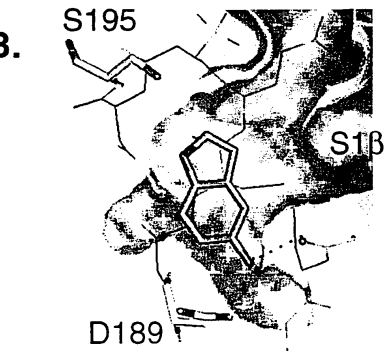
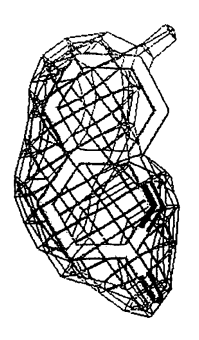


Figure 8

Α.



B.

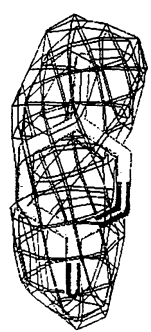
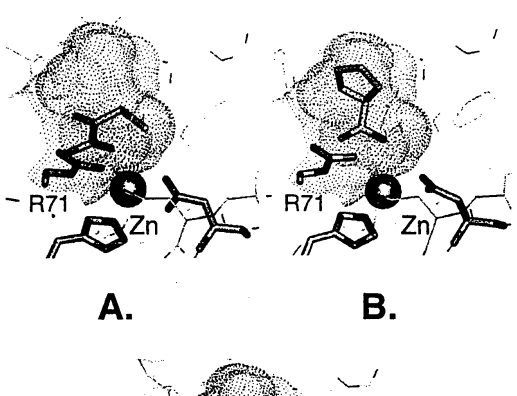
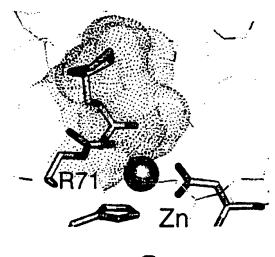


Figure 9



Figure 10





C.

Figure 11

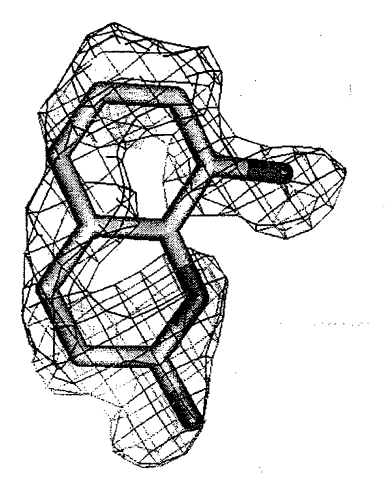


Figure 12

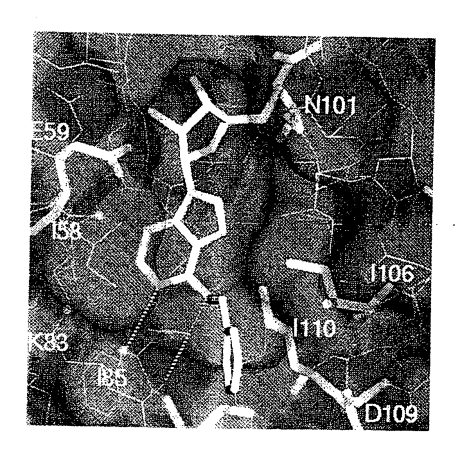


Figure 13

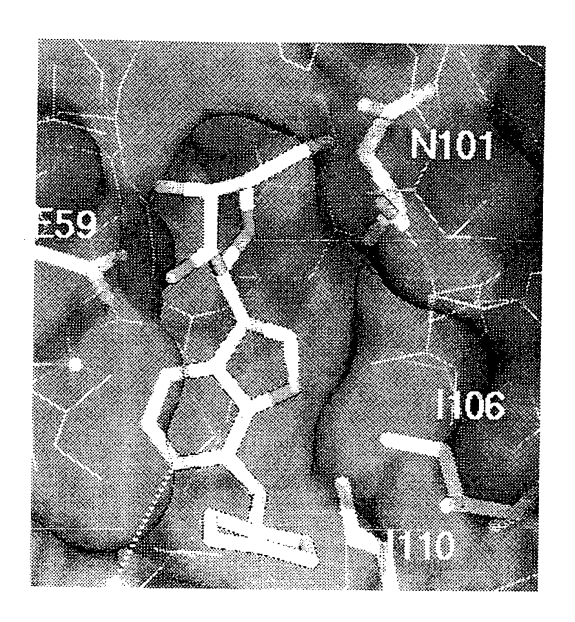


Figure 14